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- 6 Use of 5-(3,4-dimethoxyphenethyl)methylaminol-2-(3,4-dimethoxy-phenyl)-2-isopropylvaleronitrile.
- 5-((3,4-dimethoxyphenetyl)methylamino)-2-(3,4-dimethoxy-phenyl)-2-isopropylvaleronitrile or a pharmaceutically acceptable salt thereof is effective to prevent metastasis of cancer.

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Use of 5-((3,4-dimethoxyphenethyl)methylamino)-2-(3,4-dimethoxy-phenyl)-2-isopropylvaleronitrile

This invention relates to a novel agent for preventing metastasis of cancer, i.e. an antimetastatic agent.

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More particularly it relates to an antimetastatic agent comprising 5-[(3,4-dimethoxyphenethyl)methylaminol-2-(3,4-dimethoxyphenyl)-2-isopropyl-

valeronitrile of the formula

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or a salt thereof as an active ingredient.

In the recent statistics of the survey, cancer has occupied the first place in death causes in Japan, instead of cerebrovascular diseases. 24% of deaths, i.e. one among four, died of cancer. This mortal disease causes indescribable pain not only to the body but to the mind of a patient. In addition, cancer would most frequently attack those in the prime of life (i.e. in fourties to fifties) and playing important roles both in society and in their own homes so that their families also suffer from serious mental and economical damages.

Therefore various studies to reveal the fundamental cause of cancer and to establish epoch-making processes for the treatment and diagnosis thereof -3-

have been carried out all over the world to thereby gain ascendancy over cancer step by step. These studies have brought about significantly improved treatments and diagnosis of cancer, so that it can be completely cured in most cases if detected early enough.

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Even if the original tumor is completely removed by early diagnosis followed by a surgical operation, however, tumor cells would metastasize to another organ at the time of the diagnosis in more than half cases.

That is, many patients died of metastasis of cancer.

Accordingly it is one of the most important problems in the treatment of cancer to prevent its metastasis.

and important characteristic of cancer, would comprise many steps such as liberation of cancerous cells from the primary portion, transfer via blood or lymph vessels, adhesion to a blood or lymph vessel of an organ, infiltration and growth. The metastasis of cancer is an important factor governing the recuperation of a patient. However studies thereon still remain significantly backward since appropriate experimental system to evaluate the metastasis is quite limited. The mechanism of metastasis has not been clarified up and few countermeasures have been established at present.

In order to lower the mortality from cancer, it is a very important problem to prevent and treat the metastasis. Few antimetastatic agents, however, have been known to date.

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Under these circumstances, we have tried to develop an agent for preventing metastasis of cancer, i.e. an antimetastatic agent, for a long time and found that verapamil, i.e. 5-[(3,4-dimetho-xyphenethyl)methyl]-2-(3,4-dimethoxyphenyl)-2-iso-propylvaleronitrile of the following formula or its salt such as hydrochloride would be unexpectedly effective as an antimetastatic agent.

Accordingly it is an object of the present invention to provide a novel antimetastatic agent.

Verapamil of the above formula has been used as a therapeutic agent for ischemic cardiac diseases

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in treating, e.g., stenocardia, coronary arteriosclerosis (chronic ischemic cardiac dis ases, silent ischemic cardiac diseases and arteriosclerosis cardiac diseases), and myocardial infraction.

Verapamil hydrochloride has a melting point of 138.5 to 140.5°C (decomp.).

To further illustrate the present invention, the following examples will be given.

Example 1

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Effect of verapamil on plumonary metastasis of B16 melanoma BL-6

B16 melanoma BL-6, isolated by Dr. Hart et al. in U.S.A., is a cell line which infiltrates 15 through a bladder membrane and shows metastatic potential. 5 x 104 cells of B16 melanoma BL-6 were inoculated into the tail vein of a male C57BL/6J mouse. Verapamil hydrochloride was administered intraperitoneally once a day two 20 days before the inoculation of tumor cells and three days thereafter, that is, six times in total. On the 25th day of the transplantation, the mouse was anatomized to observe the metastasis to the lungs. The degree of the metas-25 tasis was evaluated by the number of plumonary nodules. The evaluated values were represented by range, median and mean + DS. a mark "a" indicates that a significant difference has been observed when compared with a control, 30 that is, p is smaller than 0.05 (Student's t-test). Ten mice were used per a group. Table 1 shows results.

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Dose of verspamil			No. of pu	No. of pulmonary nodules	ules	
hydrochloride (mg/kg)	Hedian	Range	% to control	Mean ± 8D	to control	Bignificant Difference
. 30	4.5	3~ 9	32	2.1±0.8	29	s;
40	12	1~19	96	8.8±6.8	56	6
. 09	9.5	2~13	68	8.0±4.3	46	5 5
Control	14	4~33	100	17,519.3	100	

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Example 2: Metastasis of B16 melanoma BL-6 to lungs and lymphonodi

25 x 10⁴ cells of Bl6 melanoma BL-6 were transplanted to the right forefoot of a C57BL/6J male mouse. Cancerous cells would spantaneously metastasize to the right nodi lymphatic axillares and lungs with the elapse of time. Verapamil hydrochloride was intraperitoneally administered once a day from the fifth to 16th day (i.e. 11 times) after the transplantation of the cancerous cells. On the 17th day of the transplantation, the right forefoot including the primary tumor was cut off. On the 38th day the mouse was anatomized to determine the number of plumonary nodules.

15 Table 2 shows the result.

Table 2

Dose of verapamil			No. of pu	No. of pulmonary nodules	ules	
hydrochloride (mg/kg)	Median	Range	% to control	Mean ± 8D	& to control	Significant Difference
30	11	3~21	147	12.3± 6.5	75	
40	5	a . 1∿12	67	5.8± 3.7	35	
. 20	1.5	0~12	20	3.5± 4.3	21	4 5
Control	7.5	1~42	100	16.5±17.1	100	

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Example 3

Effect of verapamil on plumonary metastasis of highly metastatic clone NL-17 of mouse colonic cancer colon 26 adenocarcinome

5 x 10⁴ cells of a highly metastatic cell
strain clone NL-17 were transplanted into a vein of
a BALB/C femal mouse. Verapamil hydrochloride was
administered intraperitoneally to the mouse once
a day two days before the inoculation of tumor cells and three
days thereafter, i.e. six times in total. On the
23th day of inoculation, the mouse was anatomized
to determine the number of metastatic pulmonary
nodules. Results are shown in Table 3.

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Dose of verapamil			No. of pu	No. of pulmonary nodules	ules	
hydrochloride (mg/kg)	Median	Range	* to control	Mean ± SD	control	Significant Difference
09	7	96~0	2.1	13.1±31.3	15.6	ed .
. 27	1.5	0.67	3.2	18.6±24.5	22.1	æ
Control.	47.5	47.5 5~>200	100	84 ±71.3	100	

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Example 4

Effect of verapamil on plumonary metastasis of highly metastatic clone NL-22 of mouse colonic cancer colon 26 adnocarcinome

1 x 10⁶ cells of a highly metastatic cell strain NL-22 of mouse colonic cancer colon 26 were transplanted into the right forefoot of a BALB/C female mouse. Cancerous cells would spontaneously metastasize to the lungs with the elapse of time.

Verapamil hydrochloride was intraperitoneally administered to the mouse once a day from the sixth to 12th day of the transplantation, i.e. six times in total. On the 13th day of the transplantation, the right forefoot including the primary carcinoma was cut off. On the 29th day, the mouse was anatomized to determine the number of plumonary nodules.

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Table 4

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	Significant Difference		đ	eg .	
ules	% to control	59	42	50	100
No. of pulmonary nodules	Mean ± SD	33.1±14.6	23.5± 8.5	27.8±21.8	55.9±31.8
No. of pu	% to control	91	59	50	100
	Range	12v 55	10~ 41	70 74	22~126
	Median	34.5	22.5	19	38
Dose of verapamil	hydrochloride (mg/kg)	50	. 09	75 .	Control

Examples 1 to 4 as shown above clearly indicate that the verapamil hydrochloride according to the present invention remarkably prevents metastasis of cancer not only in a single experimental system but also in various experimental systems for cancer metastasis in animals.

Accordingly the verapamil according to the present invention is useful as an excellent agent for preventing metastasis of cancer, i.e. an antimetastatic agent.

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The dose of the verapamil of the present invention as an antimetastatic agent depends on various factors such as the type of cancer and the condition of the patient. It may be usually administered to an adult orally or parenterally in a dose of 10 to 500 mg once to four times a day without any limitation.

It may be formulated into various forms such as powder, grain, granule, tablet, capsule and injection. Formulation may be carried out in a conventional manner with the use of conventional carriers.

In addition to the use as a therapeutic agent administered to cancerous patients, the verapamil of the present invention is further available in

preventing metastasis in those who have received medical treatments such as chemotheracy, endocrinotherapy and immunotherapy, radiotherapy or surgical treatments.

5 Needless to say, the agent of the present invention may be simultaneously administered with other carcinostatic agents.

Toxicity of the verapamil hydrochloride as used in the present invention will now be shown.

10 Acute toxicity

Table 5 shows LD₅₀ (mg/kg) thereof.

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female

Animal Sex Oral Subcutaneous Intramuscular Intravenous Mouse Male 163 68 7.6 Male 108 107 118 16 Rat Pemale 126 Male and

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Table 5

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Dog

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As described above in detail, the verapamil of the present invention is remarkably effective as an antimetastatic agent. Since metastasis is the cause of deaths due to cancer in most cases, the present invention is extremely valuable.

CLAIM:

Use of 5-((3,4-dimethoxyphenetyl)methylamino)2-(3,4-dimethoxy-phenyl)-2isopropylvaleronitrile of the formula given
below or a pharmaceutically acceptable salt
thereof for the preparing of a composition for prevention of metastasis of cancer.

CH₃O CN

CH₃O CH₂ CH₂

CH₃O CH₂ CH₂

CH₃O CH₂ CH₂

CH₃O CH₂ CH₂ CH₂

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EUROPEAN SEARCH REPORT

Application number

EP 85 10 4787

	DOCUMENTS CON	SIDERED TO BI	RELEVANT			
Category		vith indication, where appearant passages	propriate,	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. CI.4)	
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